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Increasing the reaction kinetics of alkali activated fly ash binders for stabilisation of a silty sand pavement sub-base

^{a,*} Sara Rios; ^b Nuno Cristelo; ^c Tiago Miranda; ^d Nuno Araújo; ^e Joel Oliveira; ^d Ernesto Lucas

^a *Post-Doc Research fellow, CONSTRUCT-GEO, Department of Civil Engineering, Faculty of Engineering, University of Porto, 4200-465 Porto, Portugal*

^{*} Corresponding author

E-mail address: sara.rios@fe.up.pt

^b *Assistant Professor, CQVR, Department of Engineering, University of Trás-os-Montes e Alto Douro, 5001-801 Vila Real, Portugal*

^c *Assistant Professor, ISISE, Department of Civil Engineering, University of Minho, 4800-058 Guimarães, Portugal*

^d *Assistant Professor, ISISE, Department of Civil Engineering, University of Minho, 4800-058 Guimarães, Portugal*

^e *Assistant Professor, CTAC, Department of Civil Engineering, University of Minho, 4800-058 Guimarães, Portugal*

^d *PhD student, CQVR, Department of Engineering, University of Trás-os-Montes e Alto Douro, 5001-801 Vila Real, Portugal*

Abstract

The paper addresses several options to improve the reaction kinetics of alkali activated low calcium fly ash binders for soil stabilisation in road platforms. For that purpose, an experimental program was established to assess the strength evolution, with time, of different binders, based on ash, lime, sodium chloride and alkali solutions, applied in the stabilisation of a silty sand. The tests included unconfined compression strength tests, triaxial tests and seismic wave measurements performed at different curing periods. The results were compared with a binder made of Portland cement and a commercial additive specifically designed for soil stabilization in road applications. The activated ash mixtures with lime were the most performing producing a significant increase in the reactions development and, consequently, in the strength gain rate. The sodium chloride significantly improved the lime and lime-ash mixtures, but provided only a slight improvement in the activated ash mixtures.

Keywords: soil stabilization; fly ash; lime; alkaline activation, Portland cement

Introduction

With the worldwide increase in constructed surface area, the need for local soil stabilization or improvement keeps growing. Different methods have been used for decades, and several of those are still being improved and refined, while new ones keep arising as technology progresses. According to Mitchell (1981) and van Impe (1989) the stabilisation / improvement techniques can be divided in the following groups: consolidation; grouting; mechanical, chemical and physical stabilisation; and soil reinforcement. Almost all of the mentioned groups comprise one or more techniques that can be very efficiently applied in transport infrastructures. Compaction, consolidation, grouting and soil reinforcement can be used to improve the bearing capacity and deformability properties of embankments or slopes; while mechanical (surface compaction) and chemical stabilisation are very common when dealing with superficial layers.

In particular, chemical stabilisation is very efficient when the soils available nearby cannot present an adequate performance based only on mechanical stabilisation. The use of a soil-binder combination specifically developed for subgrade layers can significantly increase its strength and stiffness, avoiding the extraction of more performing materials from distant locations ([Ingles & Metcalf, 1972](#); Little, 1995; Sherwood, 1993; Little & Nair, 2009).

The most common binders for surface stabilisation are based on Ordinary Portland cement (OPC) and/or lime (Petry & Little, 2002; Xing et al, 2009). Due to the pozzolanic properties of common clay minerals, the high amount of calcium present in both slaked and unslaked lime makes it a very effective option to decrease plasticity and increase workability, with also some significant strength and stiffness increase if dry weight percentages above 6% are used ([Cristelo et al, 2009](#); [Little et al, 2010](#); Al-Mukhtar et al, 2014; Gullu, 2015; Han & Cheng, 2015; Elkady, 2016). When dealing with silty to sandy soils, OPC is usually the most interesting option (Consoli et al, 2011a; Bahar et al, 2004; [Goodary et al, 2012](#); Rios et al., 2012, 2014). However, if a significant increase in strength is necessary in a clayey soil, the

most consensual strategy is probably to mix the soil with lime, before mixing the resulting granular material with OPC ([Horpibulsuk et al., 2006](#); [Yusuf et al, 2001](#); [Yong & Ouhadi, 2007](#); [Oza & Gundaliya, 2013](#); [Khemissa and Mahamedi, 2014](#); [Saride et al, 2013](#)). Several binders have been tested, with more or less success ([Kolias et al, 2005](#); [Basha et al 2005](#); [Kaniraj & Havanagi, 1999](#); [Kamei et al, 2013](#); [Yilmaz & Ozaydin, 2013](#); [Gurbuz et al, 2015](#)), but in almost every case OPC plays a major role in the stabilisation process.

Based on the literature review, it becomes clear that cement is the most important constituent in chemical stabilisation of soil subbases, producing significant strength and stiffness increase with little or no setbacks whatsoever. Nevertheless, the decreasing tolerance regarding the environmental concern has the production of clinker as a major target, mostly due to the high amount of CO₂ released. A value between 5% and 8% of the overall CO₂ released to the atmosphere is estimated to be originated by OPC production ([Provis & van Deventer, 2014](#); [Scrivener & Kirkpatrick, 2008](#)). Therefore, there is an ongoing research effort targeting the development of more sustainable binders ([Juenger et al., 2011](#)). In particular, the use of waste materials is highly encouraged, since it allows an increase in resource efficiency, while contributing also to enhancing the circular economy (by reducing landfilled waste).

The alkaline activation technique is particularly adequate to create binders based on residues, such as fly ash or ground granulated blast furnace slag, which constitute very effective options due to their amorphous aluminosilicate microstructure. It consists on a reaction between aluminosilicate materials and alkali or alkali-based earth substances, such as sodium (Na) or potassium (K), or an alkaline earth ion, such as calcium (Ca). The reactions can be summarized in the following sequence. First, there is the destruction, by the high hydroxyl (OH⁻) concentration in the alkaline medium, of the Si-O-Si, Al-O-Al and Al-O-Si covalent bonds present in the vitreous phase of the original semi-amorphous aluminosilicate (i.e. the precursor). The Si and Al ions are released into the solution as they become available

and; at the same time, the alkaline cations – usually Na^+ or K^+ , depending on the activator – compensate the excess negative charges associated with the modification of the aluminium coordination during the dissolution phase. The resulting products accumulate for a period of time, forming an ion-rich solution, which finally precipitates and reorganizes into more stable and ordered Si-O-Al and Si-O-Si structures. If calcium is predominant, relatively to the sodium or potassium, the dissolved Al-Si ions will diffuse from any solid surface, which favours the production of a C-S-H gel phase. Otherwise, the Si and Al ions will be able to accumulate around the nuclei points, sharing all the oxygen ions and forming a Si-O-Al and Si-O-Si three-dimensional structure (the formation of Al-O-Al is not favoured). The resulting product is an amorphous alumina-silicate gel, which evolves, with curing time and crystallization, from an Al-rich phase to a Si-rich phase. The crystallization, starting almost immediately after the precipitation, is responsible for the hardening of the gel, which eventually matures into alkaline cement, with pre-zeolite as secondary products (Fernández-Jiménez and Palomo, 2005; Fernández-Jiménez et al., 2006). The precursor should always be submitted to a previous thermal treatment, capable of inducing the loss of constituent water and the subsequent re-coordination of the aluminium and oxygen ions, transforming an originally crystalline structure into an amorphous one, more susceptible to further chemical reactions. The relative presence of calcium in the precursor and/or in the activator is very important, since the speed of the reactions is highly dependent on the type of aluminosilicate gel being formed, either N-A-S-H or C-S-H. The former needs longer periods in order to mature into a stable and reliable matrix, while the latter has curing / developing periods similar to those obtained with cement-based binders (Dombrowski et al, 2007; Garcia-Lodeiro et al., 2013).

Alkaline activation has recently started to be applied in soil stabilisation applications, using fly ash as the precursor, allowing significant strength gains (Zhang et al, 2013; Sukmak et al., 2013, 2015; Cristelo et al., 2011, 2012a, 2013; Silva et al., 2013; Rao and Acharya,

2014; Yi et al., 2015; Rios et al., 2015; Phummiphan et al., 2016), but also financial and environmental benefits, which proved very competitive when compared with lime or cement-based soil stabilizers ([Cristelo et al., 2015](#)). Other works were also recently published using alkaline activation to stabilise other types of granular materials that do not include soils such as construction and demolition waste (Mohammadinia et al., 2015; Arulrajah et al., 2016), coffee grounds ([Kua et al., 2016](#)), or recycled asphalt pavement ([Saride et al., 2016](#), Hoy et al., 2016). However, constraints related with short deadlines regarding the entry into operation of most infrastructures, and namely road and railways, often require tight construction periods, which hinders the use of alkali activated low calcium (class F) fly ash, based on the mentioned slower reaction kinetics of the N-A-S-H gel, when compared with the C-S-H gel. However, in the presence of enough calcium, the two systems are very compatible, and several studies have focused on the characterisation of their interaction and coexistence ([Garcia-Lodeiro et al., 2009, 2011](#); [Puligilla & Mondal, 2013](#); [Bui et al., 2015](#)).

The need to improve the reaction kinetics of alkali activated class F fly ash binders constituted the motivation of the present paper. A viable method would have been the use of activated class C (high calcium) fly ash, which has proved very efficient for the stabilisation of a lateritic soil for bound pavement application ([Phummiphan et al., 2016](#)). However, assuming that such high calcium fly ash is not available, the need to develop alternative procedures to increase the strength gain rate is obvious. Therefore, several mixtures were prepared to find an effective method to increase the strength gain rate of a silty-sand stabilised with low calcium, alkali activated fly ash.

Some authors ([Ghosh and Subarao, 2001](#); [Kumar et al., 2007](#); [Samaras et al., 2008](#); [Consoli et al., 2011b](#)) have reported that a significant strength increase of lime stabilised clayey soils can be achieved with the addition of fly ash, enhancing the pozzolanic reaction, which is the main responsible for the mechanical properties of the soil-lime mixture. Additionally, several authors ([Ramesh et al., 1999](#); [Narendra et al., 2003](#); [Cristelo et al., 2009](#))

have concluded that the formation of such pozzolanic reaction compounds is faster in the presence of sodium chloride. According to these authors, this can be attributed to the increased solubility of the silica and to the formation of sodium calcium silicate hydrate (N-S-C-H), more voluminous and with a higher water-holding capacity than the well-known C-S-H gel, formed in the sole presence of soil and lime. Based on these findings regarding the role of fly ash and sodium chloride on soil stabilisation, additional mixtures were considered, for comparison purposes, in which fly ash was added to soil-lime mixtures, with and without the addition of sodium chloride. The addition of sodium chloride was also used to prepare activated fly ash mixtures. Finally, a specific commercial additive for road pavement applications (recently available in the market called RoadCem®) was mixed with Portland cement, and the resulting binder was used to prepare some additional specimens. The result was regarded as a threshold value to which the mixtures developed in the present work were compared with.

In short, the following mixtures were conceptually analysed throughout the paper:

- Lime alone and with sodium chloride;
- No-activated fly ash and lime with and without sodium chloride;
- Activated ash and lime with and without sodium chloride;
- RoadCem® and Portland cement

Experimental program

Soil identification

The work presented in this paper was based on a soil from Poland, which was fully characterized at the beginning of the experimental program described below. Typical geotechnical identification tests were performed in order to determine the particle size

distribution, Atterberg's limits, Modified Proctor compaction parameters, and the California Bearing Ratio (CBR). The particle size distribution curve (Figure 1) evidences a well graded soil, with almost 50% fines – silt (27.8%) and clay (16%), which was classified as a SC-SM - silty sand according to ASTM (2011) D 2487-11. Other parameters are summarised in **Erro!**
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The minerology was examined by a PANalytical X'Pert Pro diffractometer, fitted with an X'Celerator detector and secondary monochromator. The scans covered a 2θ range of 10° to 80° , with a nominal step size of 0.017° and 100 s/step. CuK α radiation, with a wavelength of $\lambda = 1.5418 \text{ \AA}$, was used. Qualitative phase identification was made using High Score Plus software, which utilises the International Centre for Diffraction Data Powder Diffraction File database (ICDD PDF-2, Sets 1-49, 1999) as a reference. X-Ray diffraction patterns of the soil showed the presence of quartz and muscovite on the mineralogical composition.

Characterisation of the binders

The fly ash was provided by a Portuguese thermo-electric power plant and, based on ASTM C 618 (2015), was classified as Class F due to the low calcium content as it is expressed on Table 2. Hydrated lime provided by the company Lusical, Lhoist was also used. The particle size distribution of both lime and fly ash are plotted in Figure 1 together with the soil particle analysis. The activator solution used was composed by sodium hydroxide and sodium silicate. The former was supplied in pellets, with a specific gravity of 2.13 at 20°C (99 wt. %), as indicated by the supplier, which was then dissolved in water to pre-determined concentration of 12.5 molal. The sodium silicate technical sheet indicates a unit weight of 1.464 g/cm^3 at 20°C , a $\text{SiO}_2/\text{Na}_2\text{O}$ weight ratio of 2.0 (molar oxide ratio of 2.063) and a Na_2O concentration in the solution of 13.0%. A sodium silicate to sodium hydroxide ratio of 0.5 was always considered. Deionised water was used in every mixture prepared during the work presented.

Some mixtures were also prepared with Ordinary Portland cement, type CEM I-42.5R, together with a commercial soil stabiliser, specifically designed for road pavements, named RoadCem®. This product can be described as a fine grain additive, based on alkali earth metals and synthetic zeolites, complemented with a complex activator. Based on the technical sheet, this product modifies and extends the chemistry of the cement hydration process and extends the crystallization process by forming long needle crystalline structures. It is able to delay or to speed up the hydration process of cement and can thus be used as a tool to custom design the mixes of required performance. It is always used in combination with cement and/or other pozzolanic materials (Marjanovic et al., 2009).

Specimen preparation and mixture composition

Preparation of the soil included drying and de-flocculation by hand. The solids were then dry mixed for 10 min in a Hobart counter mixer, and the liquid phase was carefully added, requiring an additional 10 min mixing period. Cylindrical specimens with 70 mm in diameter and 140 mm in height were then compacted for uniaxial and triaxial compression strength tests, using static compaction according to EN 13286-53 CEN (2004a). The top and bottom of the moulds were then covered with cling film and the specimens were left for forty-eight hours before the mould could be removed. After demoulding the specimens were allowed to cure for longer in the same conditions: at $20^{\circ}\text{C} \pm 1^{\circ}\text{C}$ and $90\% \text{ RH} \pm 3\%$.

Identification of the mixtures is presented in **Erro! A origem da referência não foi encontrada..** Each component of the solid phase is defined as a percentage of the total mass of solids, and the water content as the ratio between the amount of water and the amount of solids. All mixtures were identified using the following code: ‘S’ (soil); ‘FA’ (fly ash); ‘L’ (lime); ‘C’ (sodium chloride); ‘AA’ (alkali activated); ‘OPC’ (Ordinary Portland cement) and ‘RC’ (RoadCem). Some of these letters were followed by a number, indicating the respective dosage. In the first step the specimens were moulded on the maximum dry unit weight and

optimum water content, obtained for the soil, using a modified Proctor test (**Erro! A origem da referência não foi encontrada.**), but in the second and third steps the presence of lime and the consequent cationic exchange and flocculation prevented a higher compaction degree. Therefore, in steps 2 and 3, mixtures were moulded at 88% of the optimum dry unit weight, corresponding to a maximum dry unit weight of 18.8 kN/m^3 . All specimens were moulded with a water content of 5%, corresponding to 88% value in the Proctor curve, with the exception of the geopolymeric mixtures, which required a higher water content in order to fulfil the criteria related with the activator/ash ratios. For comparison purposes, a specimen of unstabilised soil was also prepared, being compacted at 88% of the Proctor maximum value.

Uniaxial compression strength testing

An Instron® electro-mechanical load frame, fitted with a 25 kN load cell, was used for the unconfined compressive strength tests. The tests were carried out according to EN 13286-41 (CEN, 2003a) under monotonic displacement control, at a rate of 2 mm/s, and the entire stress-strain curve was obtained from each test.

Triaxial testing

Consolidated undrained triaxial compression tests (CU), following CEN ISO/TS 17892-9 (CEN, 2004b), were performed on the original soil, after isotropic effective consolidation stresses (σ'_c) of 33 kPa, 100 kPa and 300 kPa. Additionally, drained triaxial compression tests (CD) were performed in two of the bounded mixtures – S_FA15_L5_AA3 and S_FA10_L5_AA3 – previously cured for 28 days, using effective isotropic consolidation stresses of 50 kPa, 100 kPa and 200 kPa. These particular mixtures were chosen due to the fact that they reached the most performing behaviour in the UCS tests.

After the installation in the triaxial cell, water was allowed to percolate through the specimen until a volume of water higher than twice the volume of voids was obtained. This

facilitated saturation because removed the air bubbles present in the specimen and in the tubing system. Then, the saturation followed where cell and back pressures were increased simultaneously at a rate of 30 kPa per hour keeping an effective stress of 10 kPa, until cell and back pressure achieved 410 and 400 kPa respectively. This back-pressure value was kept constant during 24 h, assuring full-saturation which was verified by the B Skempton parameter which was close to 1.

The same servo-hydraulic load frame used for UCS tests, fitted with a 25 kN load cell, was used to apply the deviatoric load, under monotonic displacement control, at a rate of 0.01 mm/min. The axial deformation was measured by two Linear Displacement Transformers (LDTs) ([Goto et al., 1991](#)), while an additional LDT was installed to monitor the radial deformation as illustrated in Figure 2.

Seismic waves

Compression seismic (P) wave velocities were measured by ultrasonic non-destructive transducers, after curing periods of 12, 24 and 48 h, and then after 3, 7, 14 and 28 days in some of the specimens tested in unconfined compression as expressed in **Erro! A origem da referência não foi encontrada..** A commercially available equipment was used, consisting of axially aligned piezoelectric transducers, with a nominal frequency of 54 kHz and 30 mm in diameter; a waveform generator; an amplifier and an interval timer with a direct reading digital display, which showed the P-wave velocity (V_P) in real time. An oscilloscope would have been preferable, due to the lower definition of the pulse wave when travelling through heterogeneous materials, like soil. However, such equipment was not available. The equipment has an excitation voltage of 1000V and a resolution of 0.1 μ s. The equipment identified automatically the P wave travel time without any operator interference, and therefore no specific analysis was needed as in other advanced devices (Camacho-Tauta et al., 2015). In this case, standard EN 12504-4 (CEN, 2003b) was used as general reference, and

calibration of the transducers was automatically performed by the equipment, using a metallic rod with a known P-wave travelling time of 20.9 μ s. The measurements were taken along the longitudinal axis of the specimens, with the specimen vertically aligned and the transducers installed on opposite faces. The acoustic coupling between the transducers and the specimen was obtained using ultrasound gel, while firmly pressing the transducers against the top surfaces of the specimen. Each result presented is the average of ten consecutive readings.

Experimental results

Unconfined compressive strength

In the first step of **Erro! A origem da referência não foi encontrada.**, designed to obtain a guideline in terms of time needed for the reaction to produce meaningful UCS results, three different liquid/solids ratios were considered, by varying the liquid phase (the solids were kept at 85% S + 15% FA). Based on the results obtained, further activated mixtures were performed with an activator / ash ratio equal to 0.707 (the criteria for such value are discussed later in the text).

In the second step, the soil was mixed with different binders in order to evaluate the effect, on the reaction's kinetics, of each of the combinations proposed. Mixtures of soil-lime as well as soil-lime-fly ash (both with and without sodium chloride) were first performed with deionised water as the only constituent of the liquid phase. Then, the fly ash and fly ash-lime mixtures were prepared with the activator as the only constituent of the liquid phase.

A third and final step was then designed to reduce the mixture cost which included the reduction of the fly ash to 10% of the total solids (and consequently, a lower quantity of activator solution), while using only one lime percentage (10%) and 1% of sodium chloride. Also included in this final step were the OPC + RC mixtures.

At least 3 specimens of each condition were always moulded and tested so that the presented data corresponds to the average of three tests, improving the statistical confidence on the results. A unique curing period of 28 days was considered during the first step, while three different curing periods of 3, 7 and 28 days were considered in the second step. In the third step, L + FA specimens were tested after 7 days, while OPC + RC specimens were cured for 3, 7 and 28 days as expressed in **Erro! A origem da referência não foi encontrada.**

The results obtained in step 1 with distinct activator contents and a ash content of 15% was aimed to evaluate the effect of the Na_2O /ash ratio on UCS (Cristelo et al., 2012a). As such, activator/ash ratios of 0.943, 0.825 and 0.707 were considered, resulting in activator contents (liquid/solids ratio) of 14.1%, 12.4%, and 10.6%. Since the three mixtures presented a significant strength increase (Figure 3), the lowest activator content of 10.6% was selected for the following steps, even though the activator content of 12.4% achieved higher strength. This was due to the fact that a reduction in activator implies a reduction of the total cost of the mixture, since the activator is, by a significant margin, the most expensive component.

Figure 4 illustrates the strength results obtained in step 2, as well as the corresponding secant stiffness modulus at 50% of the maximum strength (E_{50}). This parameter is essential to characterise the material's constitutive model included in any finite element code, and therefore its calculation is extremely useful for design purposes. With a few exceptions, the mixtures with lower and higher strength values presented also lower and higher stiffness values, respectively. Mohammadinia et al. (2014) have reported a E_{50} /UCS ratio around 1000 for cement-treated construction and demolition materials but in this case the ratio is closer to 100 probably due to a finer grading of the soil.

Note the significantly higher strength, after every curing period, achieved by the mixtures with both lime and activated ash. While the mixture with activated ash achieved similar strength values to those obtained by the lime and lime-fly ash mixtures, the addition of

lime to the former resulted in a 4 times strength increase. This is a consequence of the faster reactions in C-S-H systems, relatively to N-A-S-H systems, as observed by the authors Cristelo et al. (2012b) during a previous work that compared the strength gain rate motivated by the activation of class C and class F fly ash.

Considering the existing design guidelines for roads, both the Portuguese road specifications (JAE, 1995) and the French (GTS, 2000) classification chart for soil-cement mixtures, suggest a minimum indirect tensile strength of 0.2 or 0.3 MPa, depending whether in situ or plant mixing is used. These values can be converted into UCS values of 2 and 3 MPa assuming a ITS/UCS=10%. The GTS (2000) specification also adds the need to have 1 MPa at 7 days of curing. Using these values as target for design, as expressed in Figure 4, it becomes clear that the soil-lime mixture do not fulfil these aims, while the soil-lime-ash mixtures stay slightly above these limits. For higher performances, corresponding to higher class treated layers as expressed in GTS (2000), activated ash mixtures are needed.

A final third step was conducted to address the possible reduction of the economic and financial cost of the binders, considering that most UCS values obtained were very satisfactory and could be slightly reduced without compromising its mechanical performance. This cost decrease was obtained with a reduction of the fly ash percentage to 10%, down from the step 2 overall value of 15%. Note that a reduction in fly ash content does not represent a significant economic or environmental saving, but the reduction of the alkaline solution content needed to activate the fly ash is, indeed, significant in terms of both of these aspects. Results presented in Figure 5 (after curing for 7 days) show that the mixtures with both lime and activated ash remain as the most performing combinations, even relatively to those mixtures based on both cement and RoadCem®.

Triaxial compression strength

Figures 6 and 7 show the stress-strain curves obtained from the triaxial tests performed on the

soil and selected bounded mixtures (with lime and activated ash), respectively as indicated in **Erro! A origem da referência não foi encontrada.** These soil-binder mixtures were selected based on the efficiency/cost ratio obtained from the results from the uniaxial compression tests presented above.

The soil presented a quasi-linear behaviour up to 29%, 27% or 14% of the maximum deviatoric stress, depending on the effective confining pressure applied of 33, 100 or 300 kPa, respectively. After this linear segment, the soil stiffness significantly decreases, due to yielding, reaching a zero value when the soil achieved a constant stress plateau at the 20% to 30% strain mark. **Erro! A origem da referência não foi encontrada.** presents the results of the stiffness modulus at 50% of the maximum deviatoric stress (E_{50}), a usual design parameter as explained before.

The bounded mixtures showed completely different stress-strain curves, which in this case are typical of cemented materials ([Rios et al., 2014, 2015](#)), with a peak stress at very low strain levels (between 0.15 and 0.30%), followed by a very abrupt strain softening due to bond degradation. The material stiffness does not seem to be very much affected by the effective confining pressure, indicating that, for this pressure range, the consolidation stress did not produce an increase in stiffness, but it did not damaged the cemented structure either. This behaviour, typical of cemented soils (e.g., Fernandez and Santamarina, 2001), is expressed in **Erro! A origem da referência não foi encontrada.** showing that the E_{50} stiffness modulus was very similar for the three tests on S_FA10_L5_AA3 specimens (approximately 3000 MPa) and also for the three tests on S_FA15_L5_AA3 specimens (approximately 4000 MPa). Note that the latter E_{50} value is almost 10 times higher than the values obtained with the unstabilized soil. Also very clear is the fact that the reduction in ash content (from 15% to 10%), had a strong influence on peak stress, which decreased between 30% to 40%, depending on the confining pressure.

P-wave measurements

The results obtained from the seismic wave measurements are presented in Figure 8 relating the P wave velocity (V_P) with curing time. It should be noted that the specimens are not saturated, and so the presented results do not represent the propagation velocity through the water, but through the soil grains. Consequently, the positive evolution of V_P with curing time is expected since V_P are directly related with the constrained modulus being therefore an indirect measurement of stiffness. The relative position between the V_P values obtained in each specimen is in agreement with the previous data presented in Figure 4, confirming the sensibility of this measurement.

However, the major advantage of this data is that being a non-destructive method it is possible to have several measurements through the curing period without needing further specimens. Figure 8 shows that there is a significant improvement in the first 7 days of curing and then the evolution continues at a smaller rate. This is especially relevant for alkali-activated ash mixtures and for specimens with lime and sodium chloride.

Discussion

Strength envelope of bounded and unbounded mixtures

The strength increase resulting from the mixture with lime and activated ash can be quantified in terms of the Mohr Coulomb failure criterion, namely from the resulting strength parameters ϕ' (friction angle) and c' (cohesion). The peak and ultimate strength envelopes of the soil and the selected bounded mixtures are presented in Figure 9, while the corresponding Mohr-Coulomb parameters are summarized in **Erro! A origem da referência não foi encontrada..** It is possible to observe that the peak friction angle almost doubled, while the peak cohesion intercept suffered also a significant increase, which is typical of cemented materials. At

ultimate states the strength of the bounded mixtures approaches that of the soil since the cohesion intercept reduces drastically. However, the values of the ultimate state friction angle are still significantly higher than those obtained for the unstabilized soil (peak or ultimate state). This indicates that not all the artificially created bonds could be destroyed in the test, instead forming small clusters that reduced the void volume between the soil particles.

Effect of sodium chloride addition

Figure 10 presents the Step 2 UCS related to the sodium chloride addition. The sodium chloride was particularly effective in increasing the strength development rate of the non-activated ash mixtures (S_L5 and S_FA15_L5), particularly after 28 days. The contribution of the sodium chloride to the reaction kinetics is clear and, after 28 days, the addition of 1% represents an UCS increase of 69% of the lime-based mixture and 59% of the lime and non-activated ash mixture. Regarding the activated ash mixtures (S_FA15_L5_AA3), it appears that the sodium chloride did not make a noticeable impact on strength increase, and has even hindered the short-term (3 days) behaviour.

Effect of lime percentage

Step 2 tests also allowed to observe the effect of hydrated lime on the reaction kinetics of the mixtures. Figure 11 compares the strength values obtained for the two lime percentages considered (5% and 10%), for each of the three curing periods. The results indicate that the increase in lime content affects the rate of strength development, especially between the 7th and the 28th day. An increase of 5% in lime content, after 28 days curing, represented strength increases of 40% (soil + lime + water), 56% (soil + ash + lime + water) and 22% (soil + ash + lime + activator).

Effect of ash percentage

Comparing step 2 and step 3 UCS results, after 7 days curing, it is possible to evaluate the effect of the fly ash content – 0%, 5% and 10% of the total solids weight (Figure 12). Overall, the results are in agreement with those from the triaxial tests presented before, where it was clear that the reduction in ash content (from 15% to 10%) represents a decrease in strength. This decrease is significantly more pronounced when the ash is activated, at least after 7 days curing.

On the other hand, it is interesting to note that the use of lime and 10% fly ash represents a strength decrease relatively to the use of lime alone. This is contrary to the results of Consoli et al. (2011b), that reported a significant strength increase when 12,5% and 25% ash were added to distinct lime percentages (from 3% to 9%), when stabilizing a similar soil, using very similar fly ash, in terms of chemical composition. This suggests the possibility of existing a minimum ash content in order to take full advantage of the pozzolanic effect of the ash. On the other hand, pozzolanic effects on strength require an extended curing period to be fully noticeable. It is possible that, for higher curing periods (when there is enough time to take full advantage of the added fly ash), the strength of the soil + lime + ash + water mixtures will increase linearly with the ash content. This is not noticeable at this early curing period of 7 days, where the overall particle size distribution might be prevalent in terms of UCS. Assuming that this second hypothesis is indeed correct, the gap between activated and non-activated mixtures will probably be reduced with time. Although both mixtures have ash and calcium required to develop pozzolanic reactions, in the case of the activated mixtures there is also a N-A-S-H gel type forming, which will hinder the development of calcium-based reactions after medium to long term curing periods.

Comparison of different binders

Figure 13 compares the different binders considered: lime; activated ash; lime and ash; lime and activated ash; and OPC with RoadCem. As it was already observed in previous figures, the activated ash mixed with lime always showed the best performance, both in terms of final (28 days) strength and in terms of strength gain rate. Somehow surprisingly, the OPC + RoadCem mixture showed a rather slow strength increase, when compared with the remaining mixtures. However, it should be taken in consideration that the relative dosage might have a strong impact on these results, i.e., to have comparable strength values a higher dosage of OPC and RoadCem® might be necessary. On the other hand, the OPC content was already significant (8% and 12%), indicating that for the same strength values the use of alkali-activated ash may be more economic.

Conclusions

The paper compares the performance of different binders, based mainly on lime and fly ash, with or without alkaline activation as the liquid phase. It is highlighted that the alkaline activation reaction kinetics may be improved or reduced depending on the mixture composition. Lime, ash, sodium chloride and the alkaline activator were used in different combinations and compared to Portland cement and a specific new additive, recently available in the market for road applications. The following conclusions may be taken:

- The lime + activated ash mixtures were the most performing in terms of uniaxial compression strength and axial stiffness;
- The sodium chloride significantly improved the lime and lime-ash mixtures, but provided only a slight improvement in the activated ash mixtures;
- A 5% increase (from 10% to 15% in terms of solids weight) in lime content represented an important strength increase after 28 days, for the lime-based mixtures,

for the lime and no activated ash mixtures and for the lime and activated ash mixtures. Stress-strain curves and strength envelopes obtained in triaxial tests for the best performing mixtures (with lime and activated ash) showed a typical strong cemented soil behaviour with friction angles above 50° and cohesion intercepts higher than 500kPa.

- Considering the existing design guidelines for roads, the soil-lime mixtures studied in this work did not fulfil the necessary requirements, while the soil-lime-ash mixtures stay above the minimum values even without the activator. For higher performances, corresponding to higher-class treated layers as expressed in GTS (2000), activated ash mixtures are needed.

From the results obtained it is possible to conclude that alkali activated binders, based on low-calcium fly ash, can be very competitive with cement, regarding soil stabilisation for transport infrastructures applications. If the calcium content of the original fly ash is increased using an additional component like lime, not only the strength gain rate is enhanced, but the final strength is also improved.

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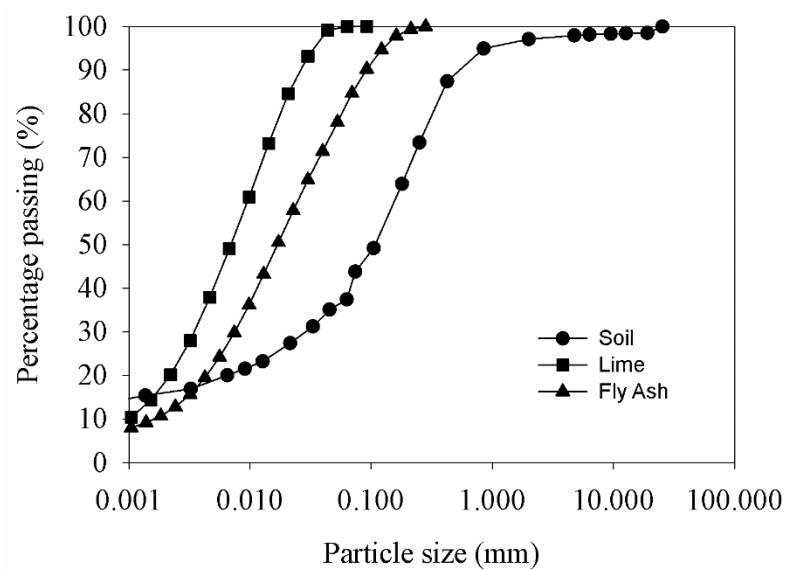


Figure 1 - Particle size distribution of the soil, lime and fly ash

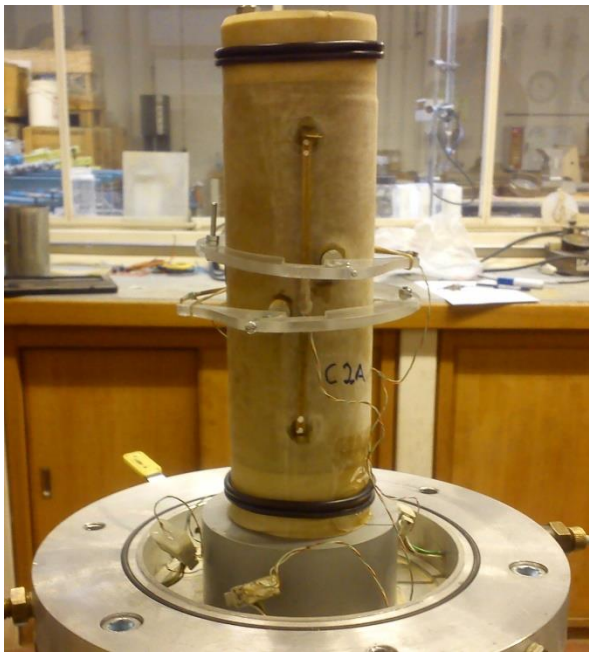
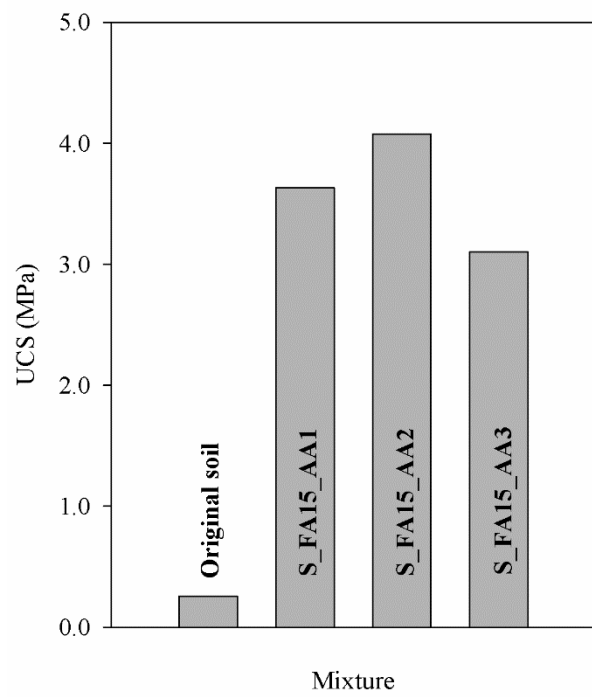


Figure 2 - Triaxial test setup showing the Linear Displacement Transformers (LDT) sensors



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740 Figure 3 - Step 1 UCS results (28 days curing)

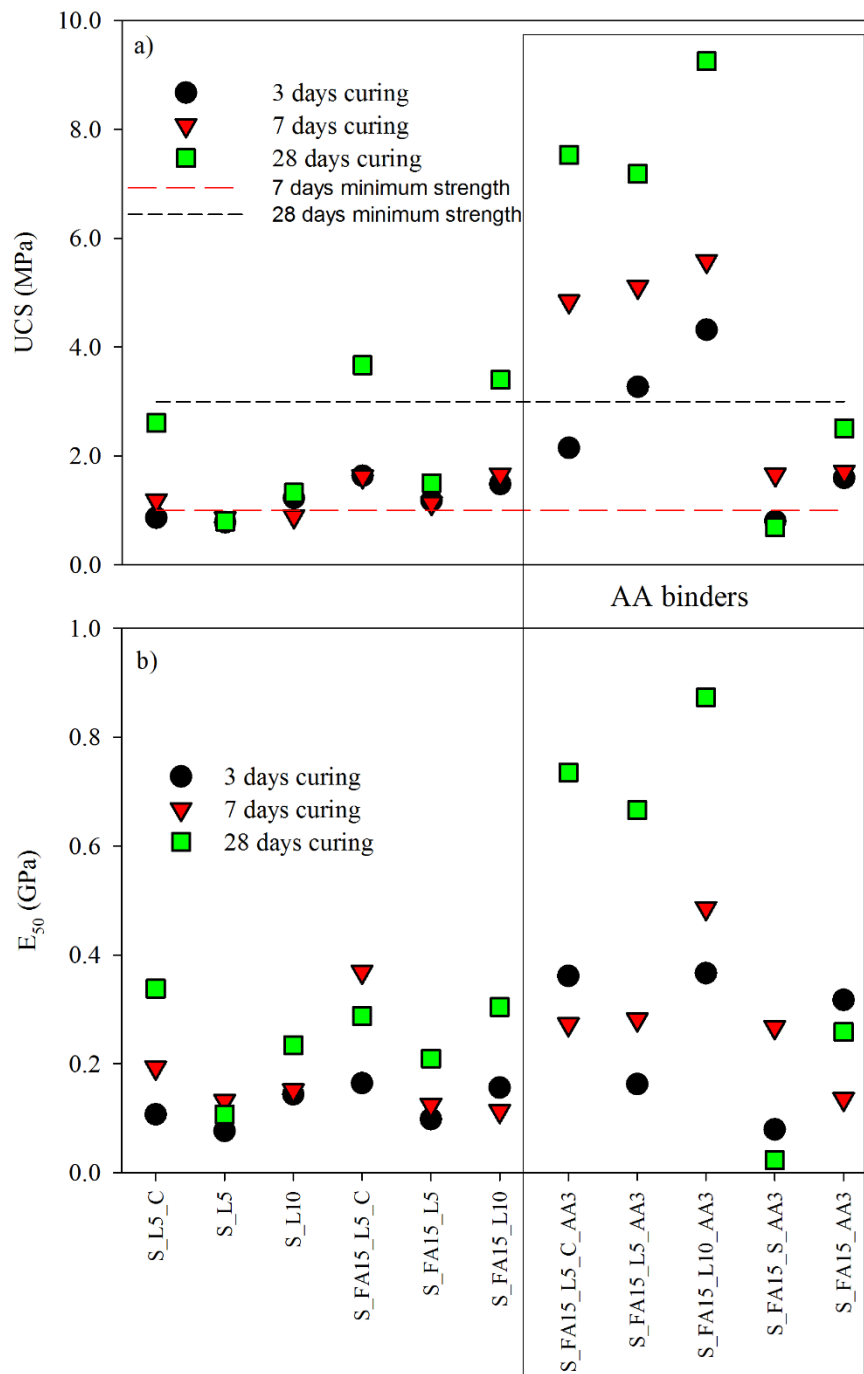
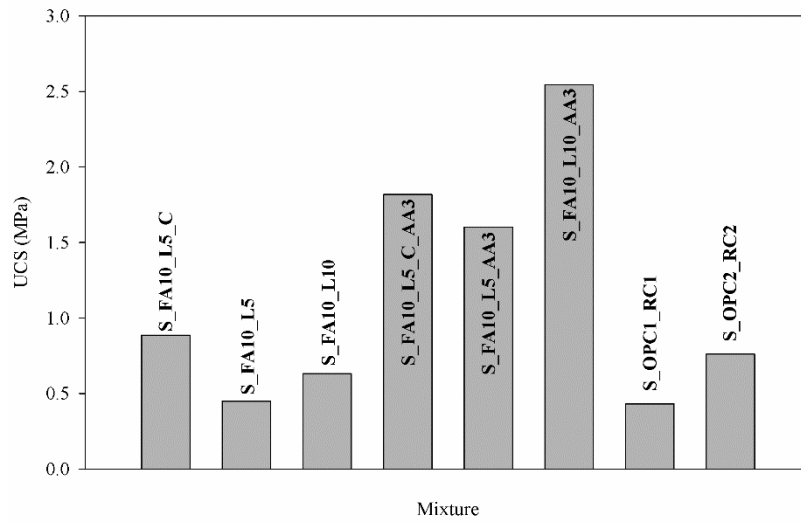
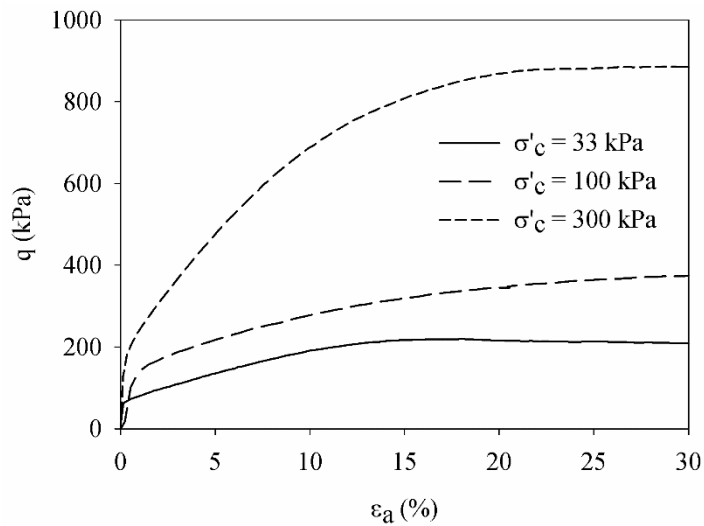


Figure 4 - UCS results for step 2 (3, 7 and 28 days curing): a) unconfined compressive strength (UCS); b) Stiffness modulus at 50% of maximum strength (E_{50}) - S (soil); FA (fly ash); L (lime); C (sodium chloride); AA (alkali activated)



745

746 Figure 5 - UCS results for step 3 (7 days curing) - S (soil); FA (fly ash); L (lime); C (sodium
747 chloride); AA (alkali activated); OPC (Portland cement); RC (RoadCem)



748

749 Figure 6 - Stress-strain curves obtained from the triaxial compression tests performed on the
750 original soil (σ'_c refers to the isotropic consolidation stress applied)

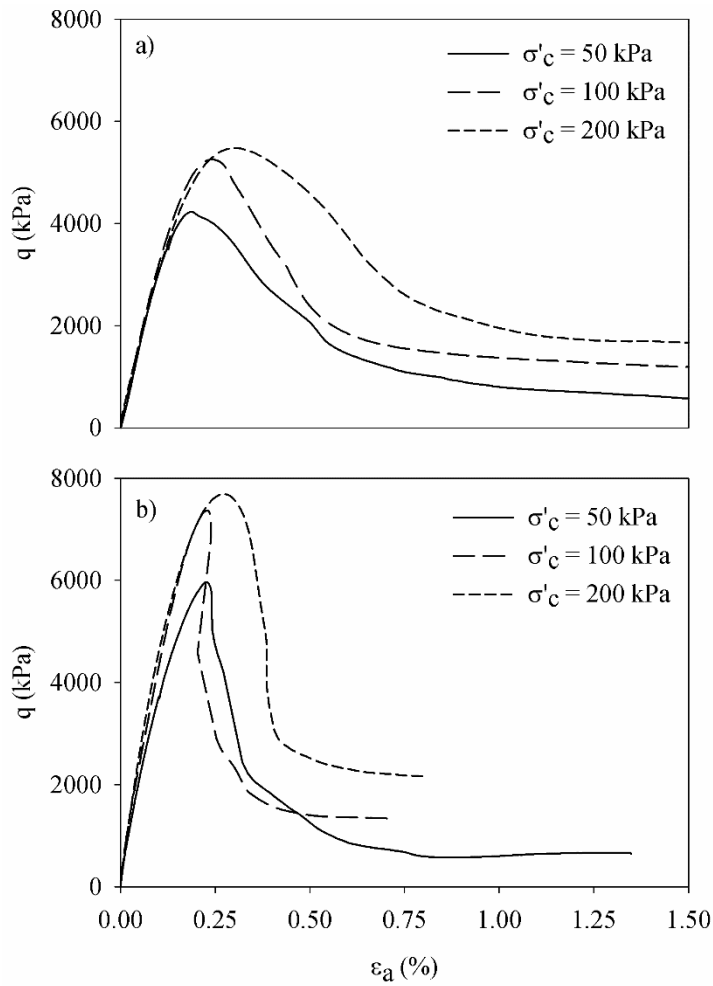


Figure 7 - Stress-strain curves obtained from the triaxial compression tests performed on the mixtures S_FA10_L5_AA3 (a) and S_FA15_L5_AA3 (b) (σ'_c refers to the isotropic consolidation stress applied)

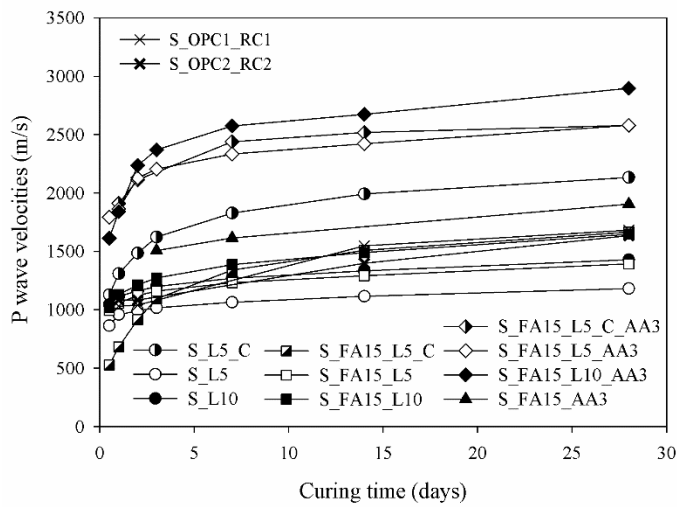


Figure 8 – P wave velocities evolution with curing time

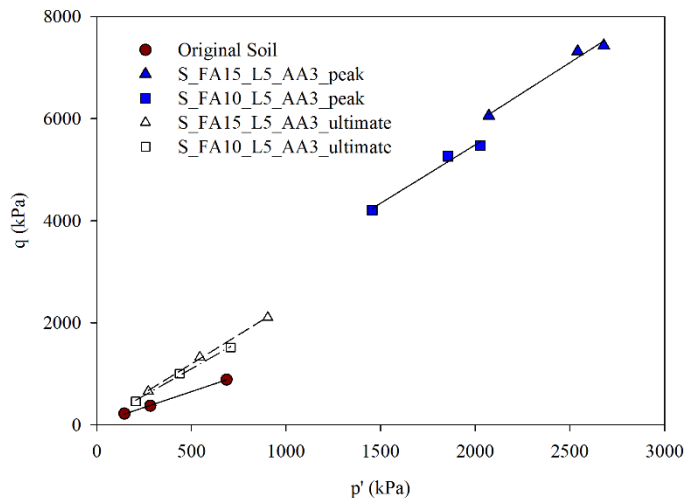


Figure 9 - Peak and ultimate state strength envelopes (the soil specimens presented very similar peak and ultimate envelopes, which are not possible to distinguish at this scale)

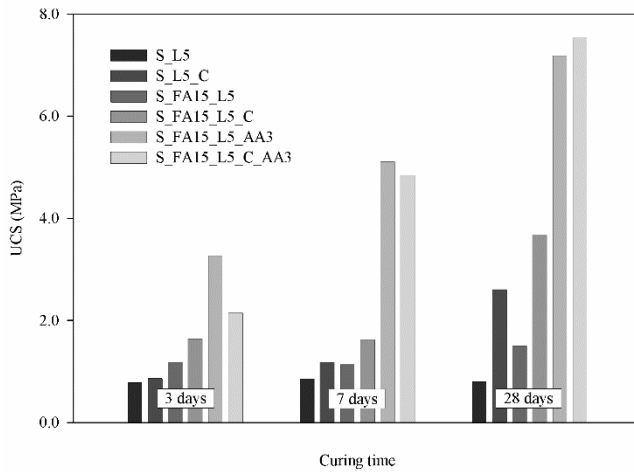


Figure 10 - Effect of sodium chloride on the uniaxial compression strength

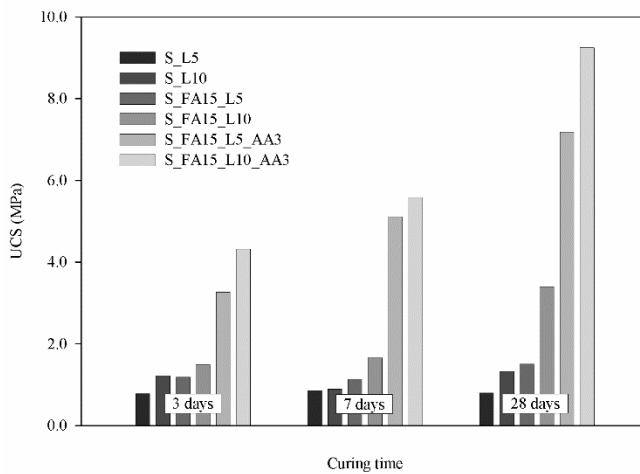


Figure 11 - Effect of lime percentage on the uniaxial compression strength

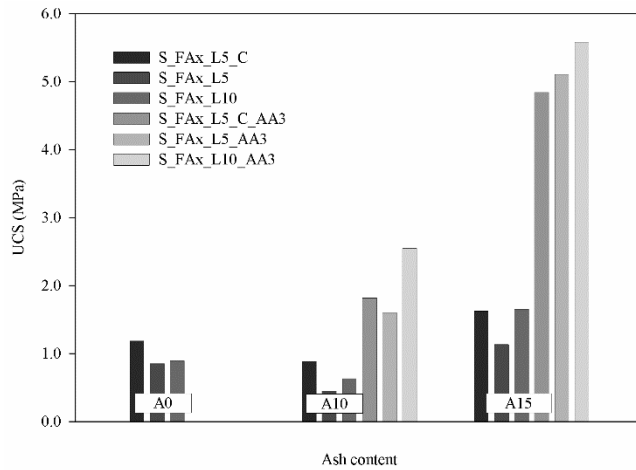


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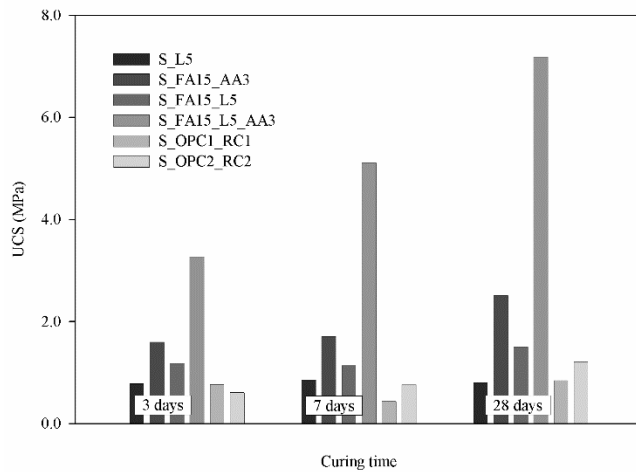


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Table 1 - Main geotechnical properties of the soil

Properties	Symbol	Value
Plastic limit (%)	w_P	13.0
Liquid limit (%)	w_L	19.5
Average diameter (mm)	D_{50}	0.105
Fines fraction (sieve N° 200) (%)	$P_{\#200}$	43.8
Uniformity coefficient	C_u	167
Curvature coefficient	C_c	6.7
Specific gravity	G	2.60
Modified Proctor test optimum water content (%)	w_{op}	8.2
Modified Proctor test maximum density (kN/m ³)	γ_d	21.29
California Bearing Ratio (%)	CBR	9
Unified Soil Classification System ASTM D 2487 (2011)	-	SC-SM silty sand

Table 2: Chemical composition of the fly ash (wt%)

Element	Si	Al	Fe	Ca	K	Ti	Mg	Na	S	P
Fly ash	48.81	21.77	14.74	3.85	4.42	1.79	1.56	1.31	1.17	0.58

Table 3 - Identification of prepared mixtures

Step	ID	Solid Phase				Liquid Phase	
		Soil (%)	Ash (%)	Lime (%)	SC(*) (%)	Activator/ Fly ash (wt)	Water content (%)
0	Soil	100	-	-	-	-	5.0
1	S_FA15_AA1	85	15	-	-	0.943	10.4
	S_FA15_AA2	85	15	-	-	0.825	9.1
	S_FA15_AA3	85	15	-	-	0.707	7.8
2	S_L5_C	94	-	5	1	-	5.0
	S_L5	95	-	5	-	-	5.0
	S_L10	90	-	10	-	-	5.0
	S_FA15_L5_C	79	15	5	1	-	5.0
	S_FA15_L5	80	15	5	-	-	5.0
	S_FA15_L10	75	15	10	-	-	5.0
	S_FA15_C_AA3	84	15	-	1	0.707	7.8
	S_FA15_AA3	85	15	-	-	0.707	7.8
	S_FA15_L5_C_AA3	79	15	5	1	0.707	7.8
	S_FA15_L5_AA3	80	15	5	-	0.707	7.8
	S_FA15_L10_AA3	75	15	10	-	0.707	7.8
3	S_FA10_L5_C	84	10	5	1	-	5.0
	S_FA10_L5	85	10	5	-	-	5.0
	S_FA10_L10	80	10	10	-	-	5.0
	S_FA10_L5_C_AA3	84	10	5	1	1.06	7.8
	S_FA10_L5_AA3	85	10	5	-	1.06	7.8
	S_FA10_L10_AA3	80	10	10	-	1.06	7.8
	S_OPC1_RC1	92	8% OPC + 0.08% RC			-	5.0
	S_OPC2_RC2	88	12% OPC + 0.11% RC			-	5.0

(*) SC stands for sodium chloride

782 Table 4 – Tests performed in each mixture and corresponding curing time

Step	ID	Curing time of the specimens tested in:		
		UCS	Triaxial tests	Seismic waves
0	Soil	0	0	
1	S_FA15_AA1	28		
	S_FA15_AA2	28		
	S_FA15_AA3	28		
2	S_L5_C	3, 7, 28		12, 24 and 48 h + 3, 7, 14 and 28 days
	S_L5	3, 7, 28		12, 24 and 48 h + 3, 7, 14 and 28 days
	S_L10	3, 7, 28		12, 24 and 48 h + 3, 7, 14 and 28 days
	S_FA15_L5_C	3, 7, 28		12, 24 and 48 h + 3, 7, 14 and 28 days
	S_FA15_L5	3, 7, 28		12, 24 and 48 h + 3, 7, 14 and 28 days
	S_FA15_L10	3, 7, 28		12, 24 and 48 h + 3, 7, 14 and 28 days
	S_FA15_C_AA3	3, 7, 28		12, 24 and 48 h + 3, 7, 14 and 28 days
	S_FA15_AA3	3, 7		12, 24 and 48 h + 3, 7, 14 and 28 days
	S_FA15_L5_C_AA3	3, 7, 28		12, 24 and 48 h + 3, 7, 14 and 28 days
	S_FA15_L5_AA3	3, 7, 28	28	12, 24 and 48 h + 3, 7, 14 and 28 days
	S_FA15_L10_AA3	3, 7, 28		12, 24 and 48 h + 3, 7, 14 and 28 days
3	S_FA10_L5_C	7		
	S_FA10_L5	7		
	S_FA10_L10	7		
	S_FA10_L5_C_AA3	7		
	S_FA10_L5_AA3	7	28	
	S_FA10_L10_AA3	7		
	S_OPC1_RC1	3, 7, 28		12, 24 and 48 h + 3, 7, 14 and 28 days
	S_OPC2_RC2	3, 7, 28		12, 24 and 48 h + 3, 7, 14 and 28 days

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784 Table 5 - Stiffness modulus at 50% of the deviatoric stress (E_{50}). Results in MPa

Mixture	Confining pressure of 33 (soil) or 50 kPa (mixtures)	Confining pressure of 100 kPa	Confining pressure of 300 (soil) or 200 kPa (mixtures)
Soil	3.68	6.84	10.21
S_FA10_L5_AA3	3194.3	3299.7	3154.3
S_FA15_L5_AA3	3901.1	4392.1	4759.9

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786 Table 6 – Mohr Coulomb failure criteria parameters

Mixture	Peak	c' (kPa)	Ultimate strength	
	ϕ' (°)		ϕ' (°)	c' (kPa)
Soil	31	15	32	0
S_FA10_L5_AA3	56	587	49	27
S_FA15_L5_AA3	58	796	56	33

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